| AD |) | |
|----|---|--|
| | | |

Award Number: DAMD17-98-1-8168

TITLE: Phage Display Breast Carcinoma cDNA Libraries: Isolation of Clones Which Specifically Bind to Membrane Glycoproteins,

Mucins, and Endothelial Cell Surface

PRINCIPAL INVESTIGATOR: Fumiichiro Yamamoto, Ph.D.

CONTRACTING ORGANIZATION: The Burnham Institute

La Jolla, California 92037

REPORT DATE: July 2001

TYPE OF REPORT: Final Addendum

PREPARED FOR: U.S. Army Medical Research and Materiel Command

Fort Detrick, Maryland 21702-5012

DISTRIBUTION STATEMENT: Approved for Public Release;

Distribution Unlimited

The views, opinions and/or findings contained in this report are those of the author(s) and should not be construed as an official Department of the Army position, policy or decision unless so designated by other documentation.

REPORT DOCUMENTATION PAGE

Form Approved OMB No. 074-0188

Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing this collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to Washington Headquarters Services, Directorate for Information Operations and Reports, 1215 Jefferson Davis Highway, Suite 1204, Arlington, VA 22202-4302, and to the Office of Management and Budget, Panegwigh Reduction Project (0704-0188). Washington, DC 20503

| 1. AGENCY USE ONLY (Leave blank | | 3. REPORT TYPE AND | | | | | |
|----------------------------------------------------------|---------------------------------|----------------------------------|----------------------------|---------------------------------|--|--|--|
| | July 2001Final Addendum (01 Jul | | | | | | |
| 4. TITLE AND SUBTITLE | anno aDNA Librariago Igal | otion of Clanca | | FUNDING NUMBERS AMD17-98-1-8168 | | | |
| Phage Display Breast Carcin | DAMDI 1-9 | 0-1-0100 | | | | | |
| Which Specifically Bind to I Endothelial Cell Surface | | | | | | | |
| 6. AUTHOR(S) | | | | | | | |
| 6. AUTHOR(S) | | | | | | | |
| Fumichiro Yamamoto, Ph | | | | | | | |
| | | | | | | | |
| 7. PERFORMING ORGANIZATION NA | AME(S) AND ADDRESS(ES) | | 8. PERFORMING ORGANIZATION | | | | |
| The Burnham Institute | AME(O) AND ADDITEOUED | | REPORT NUMBER | | | | |
| La Jolla, California 92037 | | | | | | | |
| | | | | | | | |
| E-Mail: fyamamoto@burnhan | n.org | | | | | | |
| | | | | | | | |
| 9. SPONSORING / MONITORING AC | 10. SPONSORI | RING / MONITORING | | | | | |
| U.S. Army Medical Research and | AGENCY R | EPORT NUMBER | | | | | |
| Fort Detrick, Maryland 21702-50 | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| 11. SUPPLEMENTARY NOTES | | <u> </u> | | | | | |
| 11. SOLLEWERTARI NOTES | | | | | | | |
| | | | | | | | |
| 12a. DISTRIBUTION / AVAILABILITY | | 12b. DISTRIBUTION CODE | | | | | |
| Approved for Public Rel | lease; Distribution Un. | limited | | | | | |
| | | | | | | | |
| 13. ABSTRACT (Maximum 200 Wor | edo) | | | | | | |
| 13. ABSTRACT (Maximum 200 Wor | us/ | | | | | | |
| We attempted to ident | ify phage display cDNA clo | ones that exhibited n | roteine with | carbobydrata | | | |
| | onstructed in mid copy disp | | | | | | |
| higher affinity with multi | ple number of capsid fusion | eay 17361661 10-3 ve | a saraanina | of those along the | | | |
| | | is would facilitate th | ie screening | of those clones, the | | | |
| attempts have been unsuccessful. | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| | | | | | | | |
| 14. SUBJECT TERMS | | | | 15. NUMBER OF PAGES | | | |
| Breast cancer, Phage di | 19. NOWIDER OF FAGES | | | | | | |
| Binding, carbohydrate-b | | | | | | | |
| | | | | 16. PRICE CODE | | | |
| 1 | 18. SECURITY CLASSIFICATION | 19. SECURITY CLASSIFI | CATION | 20. LIMITATION OF ABSTRACT | | | |
| OF REPORT Unclassified | OF THIS PAGE Unclassified | OF ABSTRACT Unclassifi | .ed | Unlimited | | | |

Table of Contents

| Cover | 1 |
|------------------------------|---|
| SF 298 | 2 |
| Table of Contents | 3 |
| Introduction | 4 |
| Body | 4 |
| Key Research Accomplishments | 4 |
| Reportable Outcomes | 5 |
| Conclusions | 5 |
| References | 5 |
| Appendices | |

Introduction:

In multicellular organisms, each cell is surrounded by other cells as well as a complex network of extracellular matrix. On the outer membrane, complex carbohydrate structures are present as parts of glycoproteins and glycolipids. These carbohydrate structures play an important role in interacting with proteins (lectins) (1) and other carbohydrate structures (2).

In an attempt to identify and clone unknown human proteins with carbohydrate affinity, we proposed to employ the novel technology named phage display (3). The phage display technology is based on the surface expression of the peptide sequences fused with phage capsid protein, and has been most successfully used in cloning phage particles that express variable domains of antibodies specific to certain antigens (4,5). Phage display peptide libraries made with synthetic oligonucleotides have also been utilized to identify peptide sequences that interact with a variety of bait ligands, such as proteins, peptides, DNAs, RNAs, and oligonucleotides (6,7). We have used the T7 phage cDNA display system developed by Novagen (Madison, WI) (8). Different from filamentous phage systems where the peptide sequences are fused with capsid proteins at the C-terminus, the T7 system allows the fusion of protein sequences up to 1200 amino acid residues long fused with gene 10 capsid protein at the N-terminus of proteins.

We constructed a phage display cDNA library using RNA from cells that stably expressed A transferase, and performed biopanning experiments using, as a bait ligand, crude mucin fraction containing blood group H-specific glycoproteins. Although no enrichment of the phages that expressed A transferase fusion protein was observed, selective augmentation was observed of the phages that expressed the fusion proteins with galectin-3, a soluble β -galactoside-binding (S-type) lectin (9). Because of this lectin's known affinity with the blood group-specific oligosaccharides (10,11), the results demonstrated that the phage display was useful in cloning cDNAs encoding a protein with binding capacity to carbohydrates.

Body:

During the originally proposed two years, we tried, without success, to identify cDNA clones that encode unknown proteins with carbohydrate affinity using the T7 phage system. The cDNA display libraries constructed in the T7Select 1-1 vector were primarily used in the screenings. Since the T7Select 1-1 vector displays a low copy number (0.1-1 capsid fusion per phage) of peptides or larger proteins, there was a possibility that the affinity was too weak with this system for the detection of ynknown carbohydrate-binding protein(s). Because the new vector, T7Select 10-3 vector, was developed for mid copy number display (5-15 capsid fusions per phage) of peptides and proteins, and has become available, we have repeated some of the screening experiments using the libraries constructed in this mid copy vector in the no-cost extension period. Although we hoped to identify candidate novel lectin(s) using this T7Select 10-3 vector with higher affinity, no promising candidates have been obtained.

Key Research Accomplishments:

None

Reportable Outcomes:

None

Conclusions:

Although we re-tried to identify phage clones that express fusion proteins with affinity to carbohydrate ligands using mid copy display vector, the attempts have been unsuccessful. The reason for our failure is unclear. However, different from galectins, the lectins of other family may lose the carbohydrate binding affinity by the fusion at the N-terminus. It is possible that the expression of those fusion proteins may be toxic to bacteria and the phages that express those proteins may be eliminated from the population in the library. It is also possible that those proteins are instable and may not be displayed on phage particles.

References:

- 1. Drickamer, K. Molecular structure of animal lectins. in Molecular Glycobiology (eds. Fukuda, M. & Hindsgual, O.) 53-87 (IRL Press, Oxford, UK, 1994).
- 2. Kojima, N. & Hakomori, S. Specific interaction between gangliotriaosylceramide (Gg3) and sialosyllactosylceramide (GM3) as a basis for specific cellular recognition between lymphoma and melanoma cells. J Biol Chem 264, 20159-20162. (1989).
- 3. Smith, G.P. Filamentous fusion phage: novel expression vectors that display cloned antigens on the virion surface. Science 228, 1315-1317 (1985).
- 4. Winter, G. & Milstein, C. Man-made antibodies. Nature 349, 293-299 (1991).
- 5. Clackson, T., Hoogenboom, H.R., Griffiths, A.D. & Winter, G. Making antibody fragments using phage display libraries. Nature 352, 624-628 (1991).
- 6. Wrighton, N.C. et al. Small peptides as potent mimetics of the protein hormone erythropoietin [see comments]. Science 273, 458-464 (1996).
- 7. Phage Display of Peptides and Proteins: A Laboratory Manual, (Academic Press, San Diego, 1996).
- 8. Dunn, J.J. & Studier, F.W. Complete nucleotide sequence of bacteriophage T7 DNA and the locations of T7 genetic elements. J Mol Biol 166, 477-535. (1983).
- 9. Yamamoto, M., Kominato, Y. & Yamamoto, F. Phage display cDNA cloning of protein with carbohydrate affinity. Biochem Biophys Res Commun 255, 194-199 (1999).
- 10. Abbott, W.M., Hounsell, E.F. & Feizi, T. Further studies of oligosaccharide recognition by the soluble 13 kDa lectin of bovine heart muscle. Ability to accommodate the blood-group-H and -B-related sequences. Biochem J 252, 283-287 (1988).
- 11. Sato, S. & Hughes, R.C. Binding specificity of a baby hamster kidney lectin for H type I and II chains, polylactosamine glycans, and appropriately glycosylated forms of laminin and fibronectin. J Biol Chem 267, 6983-6990 (1992).